UK Patent Application (19) GB (11) 2 156 074 A

(43) Application published 2 Oct 1985

- (21) Application No 8503900
- (22) Date of filing 15 Feb 1985
- (30) Priority data (31) 840655
- (32) 17 Feb 1984 (33) FI
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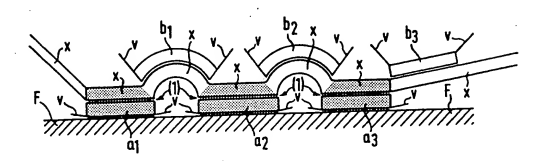
- (51) INT CL4 G01N 33/53
- (52) Domestic classification G1B 114 211 315 412 418 513 514 515 516 BT C3H 606 621 B7 U1S 1337 C3H G1B
- (56) Documents cited WO A1 8301459
- (58) Field of search G1B C3H

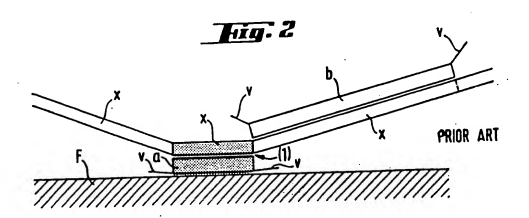
(54) Improved nucleic acid reagents and methods for their preparation

(57) The invention is related to improved nucleic acid reagents comprising arrays of nucleic acid fragments and combinations of such fragments. The preparation of such fragments by recombinant DNA techniques and their use in hybridization methods is also described.

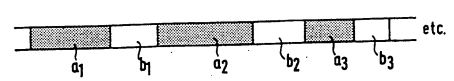
The improved nucleic acid reagents comprise two series, one labeled and one affixed to a solid carrier of at least two but preferably more arrays of alternating nucleic acid fragments, which are sufficiently homologous to sequences in the nucleic acid to be identified. Nucleic acid fragments belonging to different series must not be homologous to each other.

Hig. 1

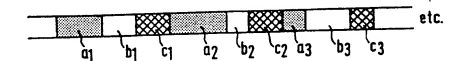


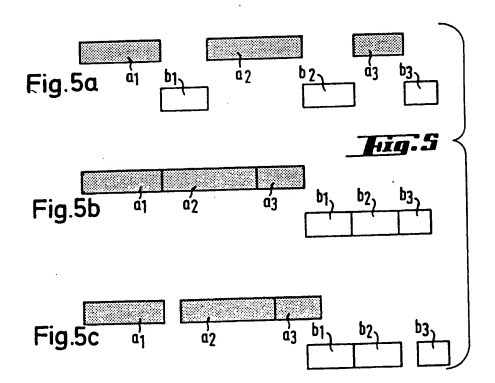


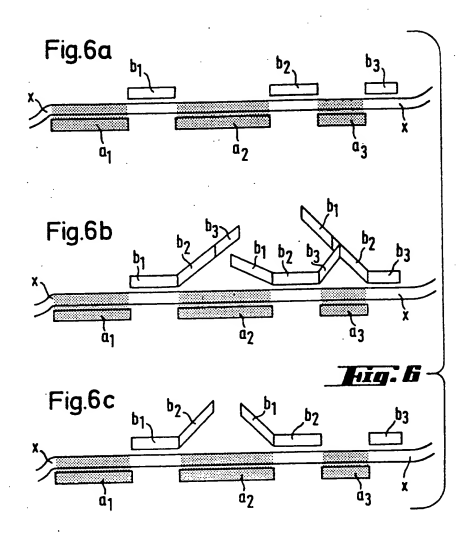




Hig. 4

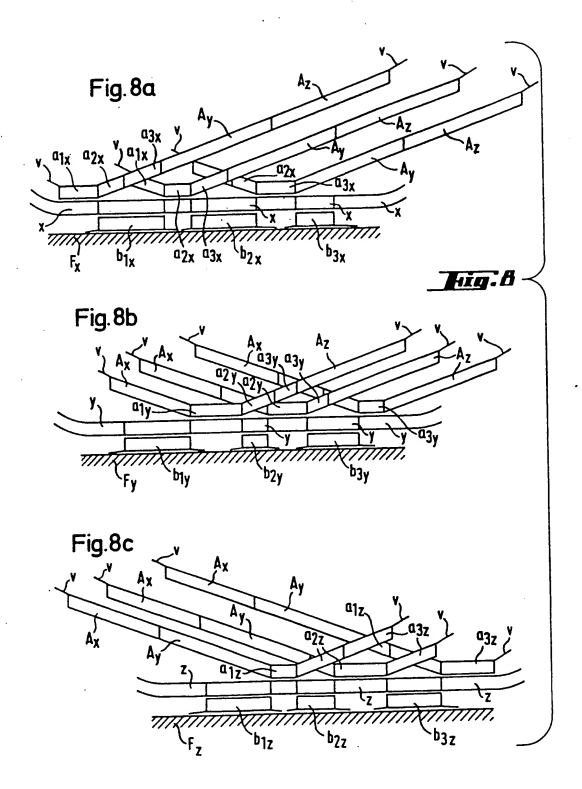






Ax Ay Az V

a_{1x} a_{2x} a_{3x} a_{1y} a_{2y} a_{3y} a_{1z} a_{2z} a_{3z}



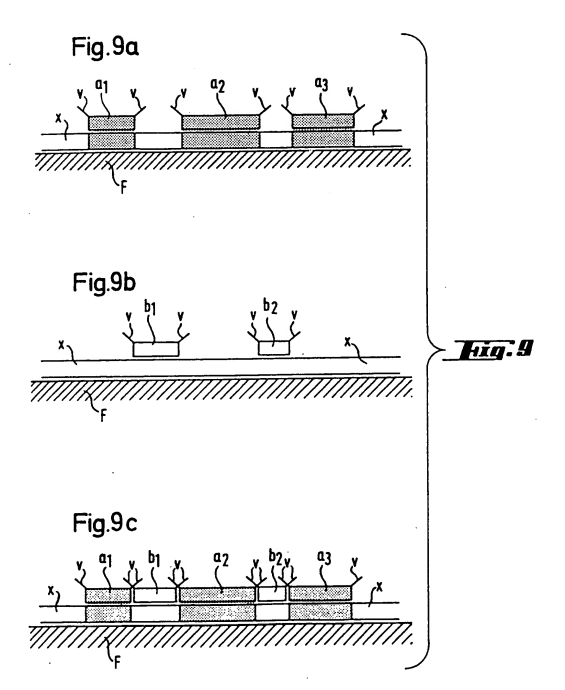


Fig: 10

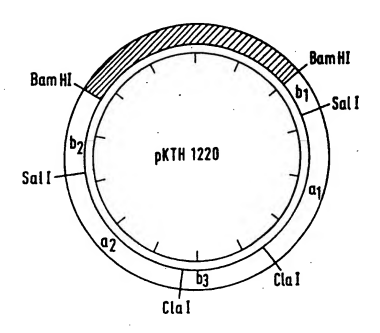


Fig.11

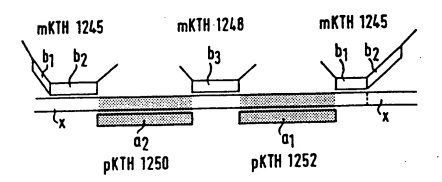
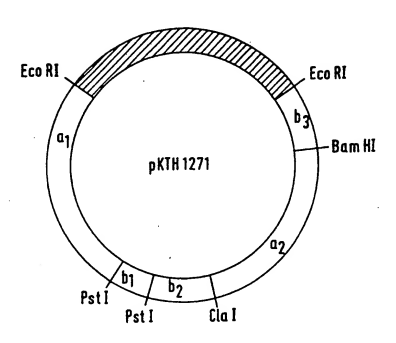
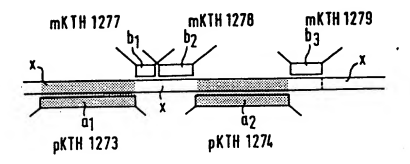


Fig.12



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SPECIFICATION

Impr ved nucl ic acid reag nts and m thods for their preparation

	Impr ved nucl ic acid reag ints and m thods for their preparation	
5	The invention relates to improved nucleic acid reagents comprising an array of nucleic acid fragments and to combinations of such improved reagents. The invention also relates to methods for the preparation of nucleic acid reagents comprised of an array of clones, and combinations of such nucleic acid reagents, by recombinant-DNA techniques, and to their use for the identification of nucleic acids by hybridization methods.	5
·10		10
15	Patent Publications Nos. 2,034,323, 2,095,833, the European Patent Publications Nos 62,286, 62,237 and 61,740), and is detected by using one labeled nucleic acid reagent which hybridizes with the nucleic acid to be identified. Other known hybridization methods include the two-step sandwich hybridization method	1.5
20	presented by Dunn and Hassell in Cell, 12, 23–36, 1977, and the one-step sandwich hybridization methods presented in the European Patent Publication No. 79,139. For the identification of the nucleic acids by the sandwich methods two separate nucleic acid reagents are needed to detect the nucleic acids present in the sample solution. One of these reagents is affixed to a solid carrier and the other is labeled. Nucleic acid reagents, both those affixed to a solid carrier and those which are labeled, are	20
25	characterized in that their base sequence is complementary, or nearly complementary, to the nucleic acid to be identified, i.e. homologous. The nucleic acid reagents used are either natural nucleic acids as such or as fragments of them. The fragments are produced, for example, by using restriction enzymes. Nucleic acid reagents have also been prepared synthetically or by recombinant-DNA techniques. Natural plasmids (US-Patent No. 4,358,535), nucleic acids from	25
	bacteriophages (US-Patent No. 4,543,535), ribosomal RNA and messenger RNA (US-Patent No. 4,302,204), or nucleic acid from different viruses (Stålhandske et al., Curr. Top.Microbiol. Virol. 104, 1983) have been used as the nucleic acid reagents. The whole virus genome has been used for identifying, for example, parts belonging to the different viruses in the messenger RNA of a hybrid virus (Dunn and Hassell, Cell, 12, 23-36, 1977). Nucleic acid reagents have	30
	also been prepared by using recombinant-DNA techniques (US-Patents Nos 4,395,486 and 4,359,535, the European Patent Application No. 79,139 and the British Patent Publication No. 2,034,323 and the European Patent Application No. 62,286). Nucleic acid reagents produced by recombinant-DNA techniques have been used either in such a way that the replicated defined DNA fragment has been purified out from the DNA of the vector, or as recombinant-DNA	35
40	molecules linked to different vectors. The previously used nucleic acid reagents produced by recombinant-DNA techniques are made up of one continuous identifying nucleic acid fragment or of several separate clones. We have developed new, more sensitive nucleic acid reagents, comprising at least two series	40
45	of alternating arrays of nucleic acid fragments prepared from either one or several segments homologous to the nucleic acid to be identified. Nucleic acid reagents which comprise such arrays of nucleic acid fragments are in sandwich hybridization tests at least twice as sensitive as the previously used nucleic acid reagents. By using the nucleic acid reagents according to the invention, or their combinations, it is possible to identify and the according to the invention and they are apposible well applies by	45
50	identify smaller amounts of nucleic acids than previously, and they are especially well applicable for sandwich hybridization methods. The higher sensitivity of the nucleic acid reagents according to the invention in sandwich hybridization methods is in part based on the fact that the use of several probes increases the quantity of labeled hybrids on the solid carrier. There may be labeled vector-derived nucleic acid along with every hybridizing probe (Figs. 1 and 2). In Figs. 1 and 2, v represents vector-derived	50
55	DNA, x the nucleic acid to be identified, b the labeled probe, a the identifying nucleic acid reagent affixed to the solid carried, and F the filter. When several probes are used, the quantity of labeled, vector-derived nucleic acid parts increases, and more label is bound to the hybrids being formed. The hybrids are thus more easily detectable. When the array of nucleic acid fragments according to the invention are used in sandwich	55
60	hybridization methods, at least two, or as sh wn in Fig. 1, three, identifying nucleic acid fragm nts ar affixed t th solid carri r. In this case th different ar as of the nucleic acid strand x to b detected may hybridize to the nucleic acid fragments affixed t the solid carri r, for example a ₁ , a ₂ , and a ₃ , at on r several points, depending on th d gre of reacti n. When the reaction reaches its final stage, a situati n according t Fig. 1 may be pr duced, in which	60
65	the sample strand forms a loop r loops to which the prob r probes, for example, b ₁ and b ₂ in	65

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5	Fig. 1, hybridize. At this time the distance of the vector-derived nucleic acid parts from the hybridization joining point (1) d creases (Fig. 1), and the hybrid is more stable than the hybrid formed by one reagent pair (prior art) shown in Fig. 2, this hybrid being of the same size as the total area of the array of nucleic acid fragments. The vector-derived parts of a hybrid formed from one reagent pair are easily broken by, for example, mechanical strain, such as shaking. In such a case the label already bound to the hybrid escapes.	5
	Since the improved nucleic acid reagents according to the invention are more sensitive than previously used nucleic acid reagents, they are suitable for demonstrating chromosomal rearrangements and hereditary diseases.	
10		10
15	The characteristics of the invention are shown in the distinguishing features of the claims, and the invention is described in greater detail in the following description and in the acompanying drawings, in which	15
	Figure 1 shows an array of sandwich hybrids, Figure 2 depicts a sandwich hybrid of the prior art, Figure 3 shows the sites of two alternating series of nucleic acid fragments in a nucleic acid	•
20	which has been selected for the preparation of an array of nucleic acid reagents according to the invention, Figure 4 shows the corresponding sites of three alternating series of arrays of nucleic acid	20
	fragments, Figure 5 shows an array of nucleic acid fragments according to Fig. 3 separate (a), joined together (b) and both separate and joined together (c),	
25	Figure 6a shows an array of sandwich hybrids which is formed when separate fragments are used,	25
30	, and position in the contract of the contract	30
	b-fragments are used. Figure 7 shows an array of nucleic acid reagents which identify different nucleic acids, Figure 8 shows an array of sandwich hybrids which are formed when the array of nucleic acid reagents according to Fig. 7, identifying different nucleic acids, are used,	
35	Figure 9 shows an array of hybrids formed by a direct hybridization method, Figure 10 shows the recombinant plasmid pKTH1220, Figure 11 shows an array of sandwich hybrids which is formed when an array of nucleic acid	35
40	fragments prepared from the recombinant plasmid pKTH1220 are used, Figure 12 shows the recombinant plasmid pKTH1271,	40
	fragments prepared from the recombinant plasmid pKTH1271 are used. Our invention relates to nucleic acid reagents composed of an array of nucleic acid fragments. These arrays of nucleic acid reagents comprise at least two, but preferably several, alternating	
45	nucleic acid fragments, up to 20 fragments, which are derived from one or several nucleic acids sufficiently homologous to the nucleic acid which is to be identified. Thereby there are obtained at least two series of alternating arrays of nucleic acid fragments, which must not be homologous to one another.	45 .
50	The arrays of nucleic acid reagents can be prepared synthetically. In this case the fragments from the two alternating series of arrays of nucleic acid fragments, must not be homologous to each other. But they must be sufficiently homologous to alternating sites in the nucleic acids to be identified. These fragments can easily be prepared by fully automatic machines after characterization of the nucleic acid sequence of the nucleic acid to be identified.	50
55	The nucleic acid reagents according to the invention are composed of separate, or joined, or both separate and joined array of nucleic acid fragments. The arrays of nucleic acid fragments may be joined to a vector, contain parts of vectors, or be	55
	totally devoid of vector parts. The nucleic acid fragments used have a minimum length of 15 nucleotides. There is no actual upper limit for length, but it is advantageous to use fragments having a length of 20–5000 nucleotides. The nucleic acid fragments are relief to the investigation of the control of	
60	nucleotides. The nucleic acid fragments acc rding to the inventinare derived either from the genome to be identified in from on part of the genominary, for example from a relatively large clonarepresenting a certain part of the genome. The arrays of nucleic acid fragments according to the invention can thus be proposed from seving a large relative property adjacent. The arrays of nucleic acid fragments thus prepared and used for	60
65	the sam reagent. The arrays f nucleic acid fragments can als b isolated from a DNA which is not id ntical t the nucleic acid t b identified but sufficiently h m I gous, so that a stable	65

hybrid is formed between the reagent and the nucleic acid to be identified. The preparation of suitable arrays of nucleic acid fragments: is by no means limited to the isolation of suitable nucleic acid fragments from the genome. There are available many equally useful methods to prepare such arrays of fragments. The man skilled in the art can pr pare arrays of nucleic acid 5 5 fragments by synthetic or semisynthetic methods. The reagents are isolated in such a way that at least two series of alternating nucleic acid fragments, a1, a2, a3, etc., and b1, b2, b3, etc., are obtained. The nucleic acid fragments belonging to the series a1, a2, a3, etc. are composed of fragments situated close to but not adjacent to one another. The nucleic acid fragments belonging to the series b1, b2, b3, etc. are . 10 also composed of nucleic acid fragments situated close to but not adjacent to one another. The 10 nucleic acid fragments belonging to the series a1, a2, a3, etc. and those belonging to the series b₁, b₂, b₃, etc. must not be homologous to each other. It is preferable that the nucleic acids belonging to the series a₁, a₂, a₃, etc. and those belonging to the series b₁, b₂, b₃, etc. are isolated in such a way that every second fragment belongs to the a-series and every second to 15 15 the b-series, as shown in Fig. 3. In Fig. 3, a₁, a₂, a₃ and b₁, b₂, b₃ are arrays of nucleic acid fragments sufficiently homologous to the nucleic acid to be identified. It is, of course, possible that even a third nucleic acid fragment series, c1, c2, c3, etc., is isolated from the same nucleic acid, as shown in Fig. 4. It is preferable that the alternating two nucleic acid reagents follow one another directly, but this is no absolute prerequisite for the invention. The nucleic acid fragment series described above can be used either as separate fragments a, 20 a2, a3, etc., and b1, b2, bc, etc. (Fig. 5a) or joined together into longer strands a1-a2-a3, etc., and $b_1-b_2-b_3$, etc. (Fig. 5b). It is, of course, possible to prepare all kinds of intermediate forms such as, for example, an a-series in which a1 is a separate fragment and a2-a3 are joined together, and in the b-series, for example, b1-b2 are joined together and b3 is separate, etc., as 25 25 shown in Fig. 5c. Fig. 6 depicts various arrays of sandwich hybrids. Fig. 6a shows an array of sandwich hybrids in which the arrays of nucleic acid fragments are separate. Fig. 6b shows an array of hybrids in which the labeled array of nucleic acid fragments are joined together. Fig. 6c depicts a case in which an array of sandwich hybrids is formed from both joined and separate labeled arrays of 30 nucleic acid fragments. In Fig. 6, x represents the nucleic acid to be identified; b₁, b₂, and b₃ 30 represent the labeled probe, and a1, a2, and a3 represent arrays of nucleic acid fragments affixed to a solid carrier. Nucleic acid fragments which belong to the b-series can, for example, be labeled in such a way that a labeled nucleic acid reagent is obtained, i.e. the probe B. The nucleic acid reagents 35 which belong to the a-series can be affixed to a solid carrier in such a way that a nucleic acid 35 reagent A bound to a solid carrier is obtained. It is, of course, alternatively possible to prepare a labeled nucleic acid reagent A, and a corresponding nucleic acid reagent B bound to a solid carrier. Such nucleic acid pairs A and B, or B and A, labeled and respectively affixed to a solid carrier 40 can be prepared for several different nucleic acids to be identified. They can be combined into 40 suitable nucleic acid reagent combinations, which are composed of different nucleic acid reagent pairs A₁ and B₁, A₂ and B₂, A₃ and B₃, etc., or B₁ and A₁, B₂ and A₂, B₃ and A₃, etc. Reagents containing arrays of nucleic acid fragments which identify different nucleic acids can also be combined so that a probe A,-A,-Z, is obtained, which, for example, comprises an array of 45 nucleic acid fragments $(a_1-a_2-a_3)_1-(a_1-a_2-a_3)_2-(a_1-a_2-a_3)_3$, as shown in Fig. 7, in which a_1 , a_2 , 45 and a_{3x} are arrays of nucleic acid fragments A, which identify nucleic acid x; a_{1y} , a_{2y} and a_{3y} are arrays of nucleic acid fragments A, which identify nucleic acid y; a12, a22 and a32 are arrays of nucleic acid fragments A, which identify nucleic acid z, and v is a vector-derived nucleic acid part. Joined arrays of nucleic acid fragments can, of course, also be used as separate fragments, 50 50 as suitable mixtures. The arrays of sandwich hybrids according to Fig. 8 are obtained by using the reagents shown in Fig. 7. If simultaneous identification of several different nucleic acids is desired, it is, of course, necessary to use separate filters, as shown in Fig. 8. Fig. 8a shows a solid carrier identifying the nucleic acid x, Fig. 8b a solid carrier identifying the nucleic acid y, and Fig. 8c a 55 solid carrier identifying the nucleic acid z. In Figs. 8a, 8b and 8c, b1, and b2, are arrays of 55 nucleic acid fragments affixed to a solid carrier and identifying the nucleic acid x; b_{1y} and b_{2y} are arrays of nucleic acid fragments affixed to a solid carrier and identifying the nucleic acid y; and b₁₂ and b₂₂ are arrays of nucleic acid fragments affixed to a solid carrier and identifying the nucleic acid z; and x, y and z ar th nucleic acids t be identified. F,, F, and F, ar the 60 respective solid carriers or filters, A,-A,-A, is a probe which identifies all the three nucleic acids 60 simultaneously, if separat solid carriers are used. The above-described nucleic acid fragment series, reagents and reagent combinations can be prepared by recombinant-DNA t chniqu s kn wn per se. A number f nucleic acid fragm nts f

different lengths ar generated, by using r stricti n enzym s, from the nucl ic acid to b 65 identified or from a part representing it. If the restriction map of the genome to be identified is

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known, it is possible to select from the genome the suitable adjacent fragments, generated by using restriction enzymes, and the fragments are isolated and amplified by using recombinant DNA techniques.

When an unknown genome is involved, an intermediate stage can be used in the preparation of the reagents, in such a way that a relatively large restriction fragment is cloned, this fragment is mapped, and the arrays of nucleic acid fragments series a_1 , a_2 , a_3 , etc., and b_1 , b_2 , b_3 etc., are produced on the basis of the information thus obtained.

It is, of course, possible to use combinations of the above methods and to use several large separate cloned restriction fragments as starting material, and to prepare several separate series, 10 which are combined to form suitable combinations.

It is advantageous to prepare the nucleic acid fragment series a₁, a₂, a₃, etc., and b₁, b₂, b₃, etc., according to the invention by using recombinant-DNA techniques in such a way that the series a is cloned into one vector, for example into the plasmid pBR322, and whereas the series b is cloned into another suitable vector, which does not have sequences in common with the previous vector. The bacteriophase M13 is an example of such a second advantageous vector. The fragments belonging to the series a can be joined to one another, and the joined series can be cloned into one vector. For example, a₁-a₂, joined together, can be cloned as a continuous insert into the same pBR322 vector. In a corresponding manner it is possible to prepare a reagent series b₁-b₂. In the cloning it is preferred to use vectors to which very large inserts of foreign DNA can be joined. For example, lambdaphage and cosmid vectors are suitable for this

Thus, two reagent pairs comprising arrays of nucleic acid fragments are needed in the sandwich hybridization method according to the invention, a reagent labeled with the label substance to be identified, i.e. a probe, and a so-called filter reagent affixed to a solid carrier.

Most commonly, radioactive isotopes are used for labeling the probes. For example in the British Patent Publication No. 2,034,323, the US-Patents Nos 4,358,535 and 4,302,204 the following isotopes are used: ³²P, ¹²⁵I, ¹³¹I and ³H. In the European Patent Publication No. 79,139, the isotope ¹²⁵I is used. Nucleic acid probes have also been modified in different ways and labeled with, e.g. fluorescent labels (French Patent Publication No. 2,518,755). Also
enzymatic or enzymatically measureable labels are used (the British Patent Publication No. 2,019,408, the European Patent Publication No. 63,879 and the French Patent Publication No. 2,519,005). The European Patent Publications Nos 70,685 and 70,687 describe a lightemitting label and labeling method, and the French Patent Publication No. 2,518,755 describes an immunologically measurable label. The lanthanide chelates described in US-Patent No.
4,374,120, especially europium, can be used as label substances. Also the biotin-avidin label

substance described by Leary et al. (PNAS 80, 4045-4049, 1983) is suitable as a label. A few examples of labels which can be used for the labeling of nucleic acid reagents according to the invention are mentioned above, but it is evident that there will be developed new, improved label substances which are also suitable for the labeling of arrays of nucleic acid fragments

40 according to the invention.

The carriers suitable for filter reagents include various nitrocellulose filters (US-Patent No. 4,358,535 and the British Patent Publication No. 2,095,833). The DDR-Patent Publication No. 148,955 describes a method of binding nucleic acids chemically to the carrier (paper). US-Patents Nos 4,359,535 and 4,302,204 describe chemically modified papers which can be used as solid carriers. Other alternatives include nylon membranes and modified nitrocellulose filters. But it is evident that there will be developed new materials which will be even more suitable for use as solid carriers according to the invention. It is, of course, possible to use also other solid carriers, such as various chromatography matrices such as triazine- or epoxy-activated cellulose, latex, etc. In principle, there are no other limitations to the selection of the solid carrier than those to be described below. It has to be possible to affix nucleic acid in a single-stranded form to the solid carrier so that these single-stranded nucleic acids can hybridize with the complementary nucleic acid. The solid carrier must also be easy to remove from the hybridization solution, or the hybridization solution must be easy to remove from the solid

The above-described combinations of the arrays of nucleic acid reagent pairs A and B, or B and A, labeled and affixed to a solid carrier respectively, and from such nucleic acid pairs made for the identification of different nucleic acids it is possible to assemble a combination A_x and B_x, A_y and B_y, A_z and B_z.

carrier. Also, the probe must not adhere to the carrier material itself so that it cannot be washed

These combinations can be used for the simultane us identification of the nucleic acids, x, y 60 and z by sandwich hybridization meth ds.

The sample is tr ated in such a way that the nucleic acids ar r l as d int th hybridization solution, and th y ar render d single-strand d. The hybridization is carried ut in a hybridization solution, to which b the nucleic acid reagents affixed t a s lid carrier and the label d 65 on s ar added. Whin hybridization has taken place, the filt rs ar lifted from the hybridization

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5	solution, if filters have been used as solid carriers. If chromatography matrices, latex, or the like have been used, the hybridization solution is removed. The solid carriers are rinsed with a suitable washing solution. The arrays of sandwich hydrids formed (Figs. 8a, 8b, 8c) are detected by methods known per se. The radioactive label is measured, for example, by autoradiography, by a scintillation counter or by a gamma-counter. For example, an enzymatic label is identified after, for example, a color reaction, by photometry or on the basis of a precipitate. Lanthanide chelates can be detected by a so-called "time resolved fluorescence" method. An immunological label is detected by immunological methods suitable for the purpose.	5
· 10	Several different mixtures can be used as the hybridization solution; the alternatives presented in the European Patent Publication No. 79,139 and US-Patent 4,302,204 are mentioned as examples. It is, of course, also possible to use other hybridization mixtures. The hybridization takes place at a temperature of 0-80°C, but is advantageous to use, for example, a temperature of 65°C. Sufficient hybridization may occur in a very short period, but it is advantageous to use	10
15	but in this case the nucleic acid reagent affixed to a solid carrier is first added to the hybridization solution. When the hybridization has taken place, the solid carrier is washed and a second hybridization is carried out in which the labeled nucleic acid reagent is present.	15
20	The above-described labeled nucleic acid reagents or reagent combination A_z , A_y , A_z , etc., and B_z , B_z , etc., can, of course, be used in direct hybridization methods. In such a case the nucleic acid sample in a solution must be divided for each nucleic acid x, y and z to be identified or, if the sample is affixed to a solid carrier, a separate sample affixed to a carrier must be prepared for each sample. The formed array of hybrids (Fig. 9) is detected by methods	20
25	known per se. In Figs. 9, F represents the solid carrier, i.e. the filter, x the nucleic acid to be identified, and v the vector-derived parts. The labeled probes used are a ₁ , a ₂ and a ₃ (Fig. 9a), b ₁ and b ₂ (Fig. 9b), and a ₁ , b ₁ , a ₂ , b ₂ ; a ₃ (Fig. 9c). As already described above, various combinations of nucleic acid reagents can be made up	25
30	from the arrays of nucleic acid fragments according to the invention. It is possible by using these combinations to identify several different nucleic acids simultaneously. Arrays of nucleic acid fragments homologous to the different nucleic acids to be identified can be used as separate fragments in the mixtures or joined together in such a manner that one probe identifying several different nucleic acids is obtained. Nucleic acid reagents affixed to a solid carrier must, of course, be kept separate in order for the identification to be successful.	30
35	Hybridization using arrays of nucleic acid fragments can be used for identifying various human, animal and plant pathogenic microorganisms. By the method it is possible to identify microorganisms present in foodstuffs, such as clostridia, salmonellae, staphylococci, which cause food poisonings. The method is suitable for the identification of contaminants present in	35
40	water, such as enterobacteria and enteroviruses. Since the sandwich hybridization test using arrays of nucleic acid fragments is a quantitative method, it is applicable to, for example, the detection and measurement of gene amplification. This characteristic is significant in, for example, the detection and treatment of cancer. The formation of a stable array of hybrids requires that the homologous sequences of the probe reagent and the filter reagent are located within a moderate, preferably less than 5 kilobase (kb),	40
45	distance from each other in the sample strand. If changes with respect to the distance between these two areas do occur, the change is cleary observable by this method. Therefore the method is also suitable for the detection of changed mRNA, chromosomal rearrangements, the rearrangement of immunoglobulin genes for expression, and hereditary diseases. It is thus	45
50	possible to construct various reagent combinations from the arrays of nucleic acid fragments. For example, for the identification of the causative agents of venereal diseases it is possible to prepare kits which include a probe which contains arrays of nucleic acid fragments which identify gonorrhea, syphilis, herpes and chlamydiae. The identification is in this case possible by using separate filters for gonorrhea, syphilis, herpes and chlamydiae.	50
55	The invention relates particular to arrays of nucleic acid fragments comprising the recombinant plasmids pKTH1220 and pKTH1271. The recombinant plasmid pKTH1220 comprises, in the plasmid vector pBR322, DNA of <i>Chlamydia trachomatis</i> L2 which is specific to the Chlamydiae. This recombinant plasmid is cloned into the host <i>Escherichia coli</i> K12 HB101. The recombinant plasmid 1271 comprises, in the plasmid vector pBR325, DNA from the cytomegal virus AD169. This rec mbinant plasmid is cloned int host <i>Escherichia coli</i> K12 HB 101. The	55
60	hosts containing the r combinant plasmids pKTH1220 and pKTH1271 have be n deposited at the culture collection Deutsche Sammlung v n Mikroorganismen (DSM), Griesebachstrasse 8, D-3400 G"ttingen, West Germany. The number of the dep sit c ntaining the recombinant plasmid pKTH1220 is DSM2825 and th number of the deposit c ntaining th r c mbinant plasmid pKTH1271 is DMS2826. The deposits will b fre ly available once the patent	60
65	application has be n mad public. The invention is described in greater detail in the following examples. Thes examples must	65

not, however, be understood as limiting the protective scope of the invention. The structure of the nucleic acid (DNA and RNA) is similar whether the question is of a nucleic acid derived from a eucaryotic or a procaryotic cell. For this reason the principles presented in the examples are equally well applicable to the nucleic acids of animals (man included), plants and microbes or viruses. Thus the reagents according to the invention can be used to detect the nucleic acids of man, animals, plants, microbes and viruses. The arrays of nucleic acid fragments can be prepared synthetically, too. The sequence of nucleic acids to be identified can be characterized and homologous arrays of fragments prepared by automatic nucleic acid preparation machines.

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10 Example 1

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(a) Arrays of nucleic acid reagents from Chlamydia trachomatis and their preparation DNA fragments suitable for the diagnostics of the Chlamydia trachomatis group were prepared from the DNA of Chlamydia trachomatis serotype L2. The DNA was isolated and fragmented by known methods, and the resulting DNA fragments were cloned into the plasmid PBR322 and 15 transferred to the host organism Escherichia coli K12 HB101, by known methods. A gene bank of the Chlamydia trachomatis L2 bacterium was obtained as a result of the cloning, i.e. a large number of recombinant plasmids, each having a separate BamHI restriction fragment of DNA derived from chlamydiae. For reagent production, recombinant plasmids containing maximally large DNA inserts derived from chlamydial DNA were selected from the gene bank. One such

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20 plasmid is the one designed pKTH1220, which has been deposited at the culture collection Deutsche Sammlung von Microorganismen under the number (DSM 2825) and the suitability of which for use as a reagent was demonstrated by a direct hybridization test. The test showed that pKTH1220 identified all of the nucleic acids derived from different Chlamydia trachomatis serotypes, but no other nucleic acids.

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The applicable fragments, obtainable by using different restriction enzymes, were selected from the pKTH1220-plasmid DNA, and some of these fragments were transferred by further cloning into pAT153 plasmid (Maniatis et al., Molecular Cloning. A Laboratory Manual, Cold String Harbor Laboratory, p.6, 1982) and some to M13 phage. Fig. 10 shows the recombinant plasmid pKTH1220, having a molecular length of 14 kb. In Fig. 10, BamHI, Sall and Clal represent the restriction enzymes used, and a₁, a₂, b₁, b₂ and b₃ illustrate the size and mutual locations of the fragments produced with the aid of these restriction enzymes. The fragments

belonging to the series b as labeled probes. Table 1 lists the sizes of the fragments and the vectors used for further cloning, the names of the recombinant plasmids, and their use.

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35 Table 1.

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		Fragment	Size	Vector	Recombinant plasmid	Use	
40	a ₁	Clal-Sall	3.0kb	pAT153	pKTH1252	Filter	40
	a ₂	Sall-Clal	2.9kb	pAT153	pKTH1250	Filter	
	b,	Sall-BamHi	0.7kb	M13mp8	mKTH1242	Labeled probe	
	b ₂	BamHI-Sali	1.4kb	M13mp8	mKTH1239	Labeled probe	
	b₃	Clal-Clal	1.7kb	M13mp8	mKTH1248	Labeled probe	
45	b1-b2	BamHI-BamHi	2.1kb	M13mp8	mKTH1245	Labeled probe	45

The fragments listed in Table 1 were isolated from an agarose gel by electroelution and were cloned into the appropriate restriction enzyme identification sites of the vectors listed in Table 1, 50 by using known methods.

50 III oresis

The fragment BamHI-BamHI 2.1kb was produced as follows: the fragments BamHI-Sall 1.4kb and Sall-BamHI 0.7kb of the plasmid pKTH1220 were separated by gel electrophoresis in agarose gel, from which they were isolated. The purified fragments were joined to each other with the aid of T4 ligase enzyme, and of the 2.1kb DNA fragments produced in the reaction, those which had free ends which were identified by the BamHI enzyme were further joined to the BamHI restriction site of the double-stranded form of the M13mp8 phage DNA. Thus there was made a recombinant phage-DNA (mKTH1245) which contains Chlamydia trachomatis DNA comprising two separate DNA fragments which are not located adjacently in the genome. However, in the genome they are I cated adjacent to the DNA reag nts pKTH1250 and 0 pKTH1252 to b affixed to the filter (Fig. 11). Fig. 11 shows an array of sandwich hybrids which is form d when the recombinant plasmids and recombinant phag s listed in Table 1 ar

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(b) Demonstration of the s nsitivity of an array of nucleic acid reagents from Chlamydia trachomatis by using the sandwich hybridization method

used as arrays f nucleic acid reag nts.

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The sensitivity of an array of nucleic acid reagents as compared with a single continuous reagent pair was studied by the sandwich hybridization method. The test was carried out using filters which all contained 10¹¹ molecules of both pKTH1250 (a₂) and pKTH1252 (a₁) DNA rendered single-stranded. The sample to be studied was the plasmid pKTH1220, which for the test was rendered single-stranded by boiling for 5 min in 0.17 M NaOH, whereafter it was transferred to 0°C and neutralized with an equimolar amount of acetic acid. The following probes labeled with ¹²⁵J, listed in Table 1, were used in the tests: mKTH1242(b₁), mKTH1239(b₂), mKTH1248(b₃) and mKTH1245(b₁-b₂).

The hybridization was performed at +65°C for 17 hours in a hybridization solution having 10 the following composition: 4 × SSC, 0.02% Ficoll, 0.02% polyvinyl pyrrolidone, 0.2% SDS, and 200 μg/ml herring sperm DNA. The filters were washed for 2 h at 50°C with a washing solution, having the following composition: 0.1 × SSC, 0.2% SDS, and were counted using a gamma-counter. The results are shown in Table 2 and are the means of five parallel tests.

15 <u>Table 2.</u>

Specimen	-		the p			
molecules/test	b 1	ъ ₂	b 3	b ₁ ,b ₂	(b ₁ -b ₂)	(b ₁ -b ₂),b ₃
 			······		· · · · · · · · · · · · · · · · · · ·	
0	37	37	33	49	39	52
10 ⁶	48	44	48	93	68	140
10 ⁷	226	236	232	396	416	686
108	1475	1415	1456	2912	2637	3580
				7		
^b 1		_			cpm/µgDNA	
^b 2		_			cpm/µgDNA	
^b 3					cpm/µgDNA	
b ₁ -b ₂	310,0	00 cpm	/test;	7 x 10'	cpm/µgDNA	
b ₁ ,b ₂	700,0	00 cpm	/test;			
(b ₁ -b ₂),b ₃	700,0	00 cpm	/test;			•

Statistically calculated, the 95% confidence limit of the tests performed without a sample (= negative controls) was regarded as the lower limit for positivity. These values were 52-54 cpm when the probe was b_1 , b_2 or b_3 , 58 cpm when the probe was b_1 , b_2 , 56 cpm when the probe was b_1-b_2 , and 65 cpm when the probe was b_1-b_2 , b_3 .

(c) Chlamydia diagnostics by using sandwich hybridization with arrays of nucleic acid fragments

Specimens taken from three men suffering from urethritis and three women suffering from cervicitis were selected for the test. *Chlamydia trachomatis* had been isolated from the male urethral specimens and the female specimens taken from the cervix. In addition, a corresponding number of similar patient specimens, from which chlamydia had not been isolated, were studied. The specimens to be examined were taken with cotton-tipped swabs which were immersed in a chlamydia sample-taking buffer containing 0.2 M saccharros, 20 mM phosphate buffer, 3% fetal calf serum, 10 μg/ml gentamicin, 100 μg/ml vancomycin, and 25 IU/ml nystatin.

Chlamydia was cultivated fr m the specim ns. The riginal specimens were also assay d by sandwich hybridizati n using an array of nucl ic acid fragm nts. The specimens were concentrated by using 2-butanol to remove liquid from them in such a way that the final volume was about 80 μ I, their concentration for the testing thus being about 3-7 fold. Thereaft r 70 mA 65 EDTA, 0.7% SDS, 200 ug/ml proteinase K enzyme were added to the specimen, and it was

treated for 15 min at 55°C and for 45 min at 37°C. Thereafter the specimen was boiled for 5 min in 0.175 M NaOH. The boiled specimen was transferred to 0°C and neutralized with an equimolar amount of acetic acid and tested. The filters and hybridization conditions described in Example 1b were used in the test. The probe used was mKTH1245 (b_1-b_2), 300,000 cpm/400 5 μ I hybridization reaction. The results are shown in Table 3.

Table 3.

Ci		
Specimen	Hybridized	Result of
	radioactivity	chlamydia culture
	· · · · · · · · · · · · · · · · · · ·	
Man 1.	151	+
Man 2.	164	+
Man 3.	154	+
Man 4.	61	-
Man 5.	76	-
Man 6.	55	-
Woman 1.	343	+
Woman 2.	509	+ .
Woman 3.	362	+
Woman 4.	57	-
Woman 5.	58	-
Woman 6.	81	-
Buffer, \overline{x}_4	30-55	
Chl. trachomatis		
L2 bacterium, 10 ⁶	419	.
	417	•
	713	
The limit for positivity in The result in Table 3 sho	the tests was 104 cpr ows that sandwich hybragnosis venereal disease	ridization using an array of nucleic acid ses. The samples which were negative in the
The limit for positivity in The result in Table 3 sho fragments is suitable for diculture tests were negative Example 2. a) An array of nucleic ac DNA fragments suitable ovirus (AD 169, ATCC VR	the tests was 104 cprows that sandwich hybragnosis venereal disease also in the sandwich hid reagents from Cyton for the diagnostics of 0–538)–(CMV). DNA was	ridization using an array of nucleic acid ses. The samples which were negative in the hybridization test. Interpretation test and their preparation by the company of the
The limit for positivity in The result in Table 3 shows the second of th	the tests was 104 cprows that sandwich hybridges also in the sandwich hid reagents from Cyton for the diagnostics of 0–538)–(CMV). DNA was by electroelution after the heir size. The eluted Dinol. The DNA thus purieened by using the Ecological sandwich hid in the cological san	ridization using an array of nucleic acid ses. The samples which were negative in the hybridization test. Inegalovirus and their preparation Cytomegalovirus were prepared from Cytomegal is isolated and fragmented by known methods or et al., J. Virol. 42, 558–582, 1982, was the EcoRI restriction fragments had been NA was extracted with phenol, whereafter it fied was joined by means of T4-ligase to the RI enzyme, and the produced recombinant-
The limit for positivity in The result in Table 3 sho tragments is suitable for disculture tests were negative Example 2. a) An array of nucleic ac DNA fragments suitable ovirus (AD 169, ATCC VR EcoRI fragment I of about solated from agarose gel be separated on the basis of the vas precipitated with ethan DNA were transferr d t Event EcoRI fragment vector op DNA were transferr d t Event EcoRI but chl ramphenic bytomegalovirus-sp cific Divirus DNA was ascertained to EcoRI-DNA fragment were	the tests was 104 cprows that sandwich hybragnosis venereal disease also in the sandwich hid reagents from Cyton for the diagnostics of C-538)-(CMV). DNA was given to be considered by electroelution after the heir size. The eluted DI hol. The DNA thus purificated by using the Ecolocoli K12 HB101 host col sensitive clines the NA insert of the correct by the Southern bill the ras derived from the DNA the DNA the summer of the correct by the Southern bill the ras derived from the DNA insert of the correct by the Southern bill the correct consistency of the C	ridization using an array of nucleic acid ses. The samples which were negative in the hybridization test. Inegalovirus and their preparation Cytomegalovirus were prepared from Cytomegal is isolated and fragmented by known methods or et al., J. Virol. 42, 558–582, 1982, was the EcoRI restriction fragments had been NA was extracted with phenol, whereafter it fied was joined by means of T4-ligase to the

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culture collection Deutsche Sammlung von Microorganismen under number DSM 2826. The recombinant plasmid was grown and purified by known techniques.

The further clonings were carried out by known techniques by using as vectors the pBR322 plasmid and the M13mp7 and M13mp8 phages. Figs. 12 shows the hybrid plasmid 5 pKTH1271 having a molecular length of about 9 kb. The array of nucleic acid fragments shown in Fig. 12 were prepared by using the restriction enzymes EcoRI, BamHI, Clal and Pstl. Fig. 12 shows the fragments obtained by using the restriction enzymes as well as their relative size and location. Table 4 lists the sizes of the fragments in question and the vectors used for the further cloning, the names of the thus obtained recombinant plasmids, and their use either as filter 10 reagents or as labeled probes. Fig. 13 shows an array of sandwich hybrids which is formed when the array of nucleic acid fragments listed in Table 4 are used.

Table 4.

15	Restriction	fragment	Vector	Designation	use	15
a ₁ a ₂ b ₁ 20 b ₂ b ₃	EcoRI-PstI ClaI-BamHI PstI-PstI PstI-ClaI BamHO-EcoRI	(3.3kb) (3.0kb) (0.6kb) (1.0kb) (1.0kb)	pBR322 pBR322 M13mp7 M13mp8 M13mp8	pKTH1273 pKTF1274 mKTH1277 mKTH1278 mKTH1279	Filter Filter Labeled probe Labeled probe Labeled probe	20

(b) Demonstration of the sensitivity of an array nucleic acid reagents from cytomegalovirus by 25 the sandwich hybridization method

The sensitivity of an array of nucleic acid reagents as compared with one continuous reagent pair was assayed by the sandwich hybridization method. The specimen in the tests was CMV DNA, which was boild in 0.17 M NaOH for 5 min. and was thereafter neutralized as in example 1b. Filters which all contained 1011 molecules of both pKTH1273(a1) DNA and pKTH1274(a2) 30 DNA, rendered single-stranded, and the following probes labeled with 125 J listed in Table 4: mKTH1277(b₁), mKTH1278(b₂) and mKTH1279(b₃) were used in the test. The probes each contained 10s cpm/µg DNA. The hybridization was carried out as described in Example 1b. The

results are shown in Table 5.

35	Table 5.							35
40	Specimen		Hyb:	40				
	molecules	:/test	b ₁	b ₂	b ₃	b ₁ ,b ₂	b ₁ ,b ₂ ,b ₃	
45				-				. 45
	0		35	33	38	45	53	
	106		38	44	46	95	125	
	4x10 ⁶		85	135	142	. 205	292	50
50	1.6x10 ⁷		203	254	265	415	645	
55	bl	310.000	cpm/	'test				55
	ъ2	320.000	cpm/	'test				
	ъ3	300.000	cpm/	'test				
~~	b1,b2	300.000	cpm	of ea	ch/te	st		60
60	b1,b2,b3	300.000	cpm	of ea	ch/te	est		

as individual nucleic acid reagents.

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- or b₃, 59 cpm when the probe was b₁, b₂, and 63 cpm when the probe was b₁, b₂, b₃.

 The results in Table 5 show that sandwich hybridization in which an individual probe reagent in used (b₁, b₂ or b₃) detects in each case 4 × 10⁶ CMV—DNA molecules. On the other hand, hybridization with a reagent of b₁, b₂ or b₁, b₂, b₃ detects as few as 10⁶ molecules of

 CMV-DNA. The results show that the array of nucleic acid reagents are four times as sensitive
- (c) CMV diagnostics by using sandwich hybridization with an array of nucleic acid reagents
 Clinical specimens were assayed by using sandwich hybridization with an array of reagents.
 10 These samples included two urine specimens from children under 1 year. These children were suspected of suffering from congenital cytomegalo disease. A lung biopsy specimen from a patient with CMV pulmonary infection was also assayed by the present sandwich hybridization.
 Both cytomegalovirus-infected and uninfected human fetal cells were also used as specimens in the test.
 - 5 A solution which contained 1% sarcosyl and 5 mA EDTA and 200 μg calf thymus DNA was added to a 10 ml urine specimen, whereafter the DNA released from the virus, together with the carrier, was precipitated using 10 ml isopropanol at room temperature. The DNA precipitate was dissolved in 200 μl of TE buffer and was brought to a single-stranded form by boiling it for 5 min, whereafter the DNA solution was cooled to 0°C and added to the hybridization solution.
- The lung biopsy specimen (a few mm³) was minced mechanically, with a knife, 200 µl of TE buffer containing 1% SDS solution and 1 mg/ml of proteinase K-enzyme was added to it. A digestion was carried out at + 37°C for 1 h whereafter the specimen was drawn into an injection syringe twice through a thin hypodermical needle. The specimen thus homogenized was boiled, whereafter it was added to the test solution.
- The cells infected with cytomegalovirus and the uninfected cells were broken up by an SDS, proteinase K treatment, homogenized and boiled, as above.

The reagents in the hybridization test were pKTH1273(a₁) and pKTH1274(a₂) on filters and mKTH1277(b₁), mKTH1278(b₂), mKTH1279(b₃) as probes, each 200.000 cpm/reaction. In other respects the hybridization, the washing of the filters and the counting of the results were 30 carried out as described in Example 1b.

The results of the present hybridization are shown in Table 6.

Table 6.

			·
35	Specimen	Hybridized radioactivity	Virus isolation
	Infected cells (10 ⁵)	3521	Not done
	Urine 1(10 ml)	243	CMV
40	Urine 2(10 ml)	3215	CMV
	Urine from a healthy		
	person (10 ml)	52	Not done
	Lung biopsy specimen	535	CMV
	Control cells 105	68	Not done
45	No specimen	65	Not done

The results in Table 6 show that it is possible, by using an array of nucleic acid reagents, to demonstrate CMV in different clinical specimens such as urine, lung biopsy specimens and cells.

The test is specific to cytomegalovirus; it does not identify human DNA, i.e. the test is not interfered by the human DNA present in the sample. In fact the type of specimen does not interfere with the specifity of test in any way.

CLAIMS

- 55 1. Nucleic acid reagents, characterized in that they comprise arrays of alternating nucleic acid fragments.
- Nucleic acid reagents according to claim 1, characterized in that they comprise two or more series of at least two but preferably more arrays of alternating nucleic acid fragments sufficiently homologous to the nucleic acid which is to be identified but not h mologous to one 60 another.
 - 3. Nucleic acid reagents according to claims 1 and 2, charact rized in that they comprise ith r s parate r joined arrays of alternating nucleic acid fragments.
 - 4. Nucleic acid reagents according to claims 1, 2 or 3 characterized in that they comprise arrays of nucleic acid fragments which either have or do not hav vector-deriv d parts.
- 65 5. Nucl ic acid reagents according to claim 1, 2, 3 or 4 characterized in that they comprise 65

	labeled arrays of nucleic acid fragments.	
	6. Nucleic acid reagents according to claims 1, 2, 3 or 4 characterized in that they comprise	
	arrays of nucleic acid fragments affixed to a solid carrier. 7. Nucleic acid reagents according to claims 1, 2, 3 or 4 characterized in that they comprise	
=	the recombinant plasmid pKTH1220 or derivatives thereof and which recombinant plasmid	5
o	contains the DNA of Chlamydia trachomatis L2 bacterium and is cloned into the host Escherichia	
	coli K12 HB101, and the deposit number of this host containing the recombinant plasmid	
	pKTH1220 is DSM 2825.	
	8. Nucleic acid reagents according to claims 1, 2, 3, 4, 5 or 6 characterized in that they	
10	comprise the recombinant plasmid pKTH1271 or derivatives thereof and which recombinant	10
	plasmid contains the DNA of Cytomegalovirus AD169 and is cloned into the host Escherichia	
	coli K12 HB101, and the deposit number of this host containing the recombinant plasmid	
	pKTH1271 is DSM 2826.	
	9. The use of nucleic acid reagents according to claims 1, 2, 3, 4, 5, 6, 7 or 8 for the	4-
15	identification of several different nucleic acids, characterized in that suitable combinations of	15
	nucleic acid reagents are assembled from arrays of nucleic acid fragments sufficiently homolo-	
	gous to these different nucleic acids. 10. The use of the nucleic acid reagents according to claims 1, 2, 3, 4, 5, 6, 7 or 8 in	
	hybridization methods, characterized in that the arrays of hybrids formed in the hybridization	
20	methods are demonstrated by methods known per se.	20
20	11. The use of nucleic acid reagents according to claims 1, 2, 3, 4, 5, 6, 7 or 8 in	
	sandwich hybridization methods, characterized in that the arrays of sandwich hybrids formed in	
	the sandwich hybrid methods are demonstrated by methods known per se.	
	12. A method for the preparation of nucleic acid reagents according to claims 1, 2, 3, 4, 5,	
25	6, 7 or 8, characterized in that the arrays of nucleic acid fragments are prepared by	25
	recombinant-DNA techniques, synthetically or semisynthetically.	
	13. A method according to claim 12, characterized in that the preparation of the arrays of	
	nucleic acid fragments comprises:	
20	(a) the isolation of a selected nucleic acids of suitable length (b) the cloning of the selected nucleic acid into suitable vectors	30
30	(b) the cloning of the selected nucleic acid into suitable vectors(c) the fragmentation of the nucleic acids by using a restriction enzymes	-
	(d) the combination of the suitable arrays of fragments into series by using suitable ligases	
	(e) the cloning of the arrays of fragments into suitable vectors, preferably fragments	
	belonging to different series into different vectors	
35		35
	(g) the fixation to a solid carrier of the either separate or joined nucleic acid fragments	
	belonging to the other series.	
	14. A method for the preparation of a nucleic acid reagent as claimed in claim 1, carried out	
40	substantially as hereinbefore described or exemplified. 15. A nucleic acid reagent as claimed in claim 1 and substantially as hereinbefore described.	40
40	15. A flucier acid reagent as claimed in claim 1 and substantially as nevemberore described.	40